Simultaneous Removal of H2S and NH3 in Coal Gasification Processes

CONTRACT INFORMATION

DOE/MT/93005--T5

Contact Number:

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OBJECTIVES

The objective of this study is to develop advanced high-temperature coal gas desulfurization mixed-metal oxide sorbents with stable ammonia decomposition materials at 550-800° C (1022-1472° F). The specific objectives of the project are to:

- (I) Develop a combined sorbent-catalyst materials shall be capable of removing hydrogen sulfide to less than 20 ppmv and ammonia by at least 90 percent.
- (ii) Carry out comparative fixed-bed studies of absorption and regeneration with various formulations of sorbent-catalyst systems and select most promising sorbent-catalyst type.
- (iii) Conduct long-term (at least 30 cycles) durability and chemical reactivity in the fixed-bed with the superior sorbent-catalyst.

BACKGROUND INFORMATION

Nitrogen (N₂) occurs in coal in the form of tightly bound organic ring compounds, typically at levels of 1 to 2 wt% on a dry-ash-free basis. During, coal gasification, this fuel-

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bound nitrogen is released principally as ammonia and nitrogen, with smaller levels of HCN. The formation of NH₃ in a coal gasification processes is a function of the fuel gas composition and the gasifier operating conditions. During the use of coal gas to generate electricity in gas-fired turbines, fuel bound(N₂) is converted to nitrogen oxides(NO_x), which are difficult to remove and are highly undesirable as atmospheric pollutants. Recent results indicate that while the efficiency of molten carbonate fuel cell (MCFC) anodes is not effected by exposure to NH₃, NO_x is generated during combustion of the anode exhaust gas. Thus, NH₃ must be removed from the coal gas before it is used in IGCC or MCFC applications.

The product stream from a high temperature, oxygen-blown gasifier, such as Texaco, contains about 2000 ppmv of NH₃, where higher concentrations (about 5000 ppmv) occur when the gasification is conducted at lower temperatures, such as in the Lurgi or GE air-blown gasifier. A range of 1500 to 3000 ppmv is considered for this study.

Removal of H₂S using zinc-based sorbents, particularly zinc titanate, to < 20 ppmv levels has been well established (Lew et al., 1989; Jothimurugesan and Harrison, 1990; Woods et al., 1990; Gupta and Gangwal, 1993,). Previous literature study indicated that catalyst have high activities for NH₃ decomposition (Krishnan et al., 1988). If desulfurization sorbents such as zinc titanate could be used along with the NH₃ decomposition catalysts to decompose ammonia present in hot coal gas, then the number of unit processes necessary to clean hot coal gas could be reduced by one.

The objective of this project is to develop successful combination of an NH₃ decomposition catalyst with the mixed-metal oxide sorbent so that the sorbent-catalyst activity remains stable for NH₃ decomposition in addition to H₂S removal under cyclic sulfidation-regeneration conditions in the temperature range of 550-800° C (1022-1472° F) and pressures up to 20 atm.

PROJECT DESCRIPTION/RESULTS AND ACCOMPLISHMENTS

The project consists of three major experimental tasks (Tasks 1-3) addressing the contract objectives described above.

Task 1: Sorbent - Catalyst Preparation and Characterization

Task 2: Experimental Testing

Task 3. Cyclic Testing

Task 2: Experimental Testing

The activities of the HART 48 and 49 sorbent-catalysts containing Mo were tested using the simulated coal gas. The H₂S removal ability is shown in Figures 3 & 4 and its ammonia decomposition activity is shown in Figures 1&2. Figures 3 & 4 shows the H₂S breakthrough profiles as a function of time. The pre-breakthrough H₂S level was below 60 ppm. Nearly complete sorbent conversion (100%) was observed at breakthrough.

The HART 48 & 49 sorbent-catalysts showed moderate catalytic activity (50-55% average conversion) for ammonia decomposition as shown in Figures 1&2. Initially, up to 150 min all the sorbent-catalysts showed a very high activity(>90%). The decline in activity after 150 min, is mainly due to the poisioning of the catalyst by H₂S.

As shown in Figure 2, in absence of steam, the sorbent-catalysts (HART-49) showed a very high activity (>95% ammonia decomposition) until breakthrough (400 min) at 700°C. As the temperature is decreased from 700 to 550°C, the activity is >90% upto about 140 min then the activity slowly declines. A marked decrease in catalytic activity is noted when steam is present. In presence of steam, at 700°C the activity is >90% upto 200 min (see Cycle No.8) then slowly declines. It seems that the steam is more deleterious than H₂S and the steam is disintegrating the catalyst support.

FUTURE WORK

As of today, HART-49 sorbent-catalysts shows very promising catalyst activity. Multicycle runs will be conducted using HART-49 sorbent-catalysts.

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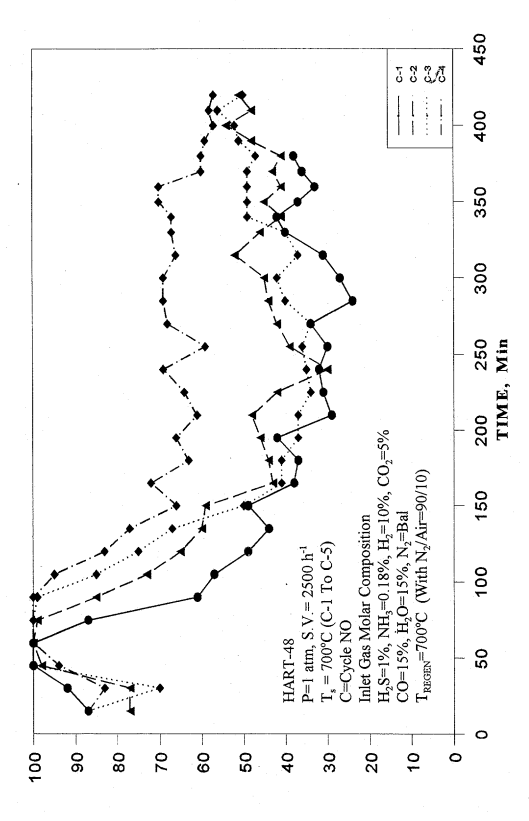


Figure 1. Conversion of ammonia on HART-48 catalyst-sorbents.

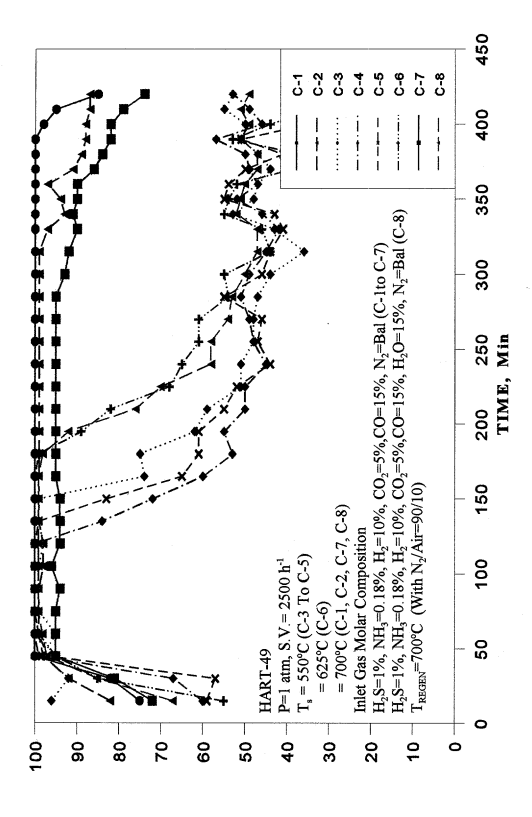


Figure 2. Conversion of ammonia on HART-49 catalyst-sorbents.

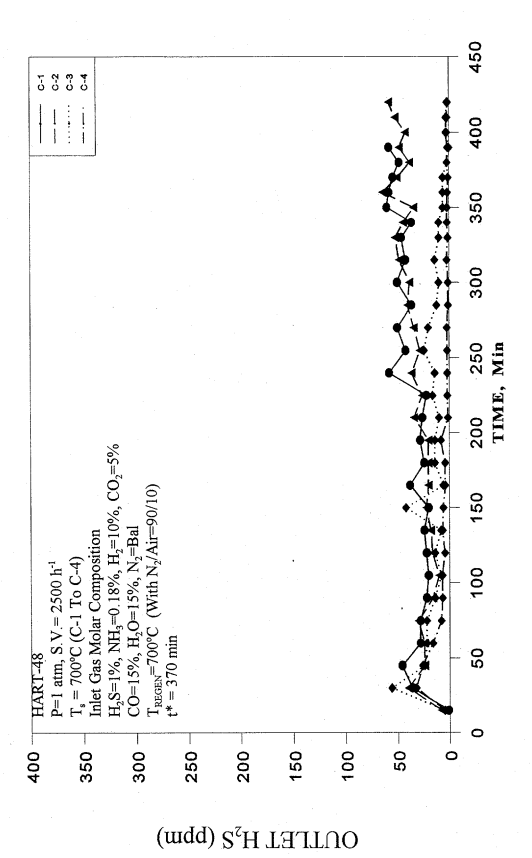


Figure 3. H₂S Breakthrough Curves in Successive Sulfidation Cycles of Sorbent-Catalyst HART-48

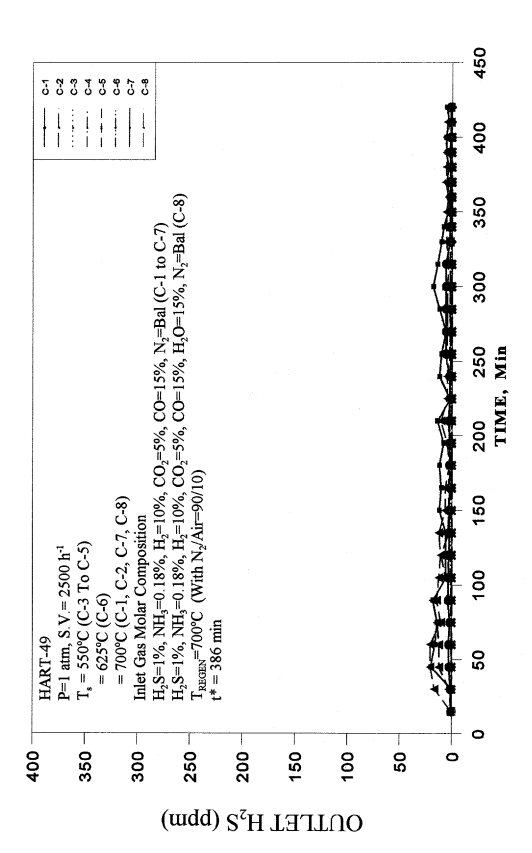


Figure 4. H₂S Breakthrough Curves in Successive Sulfidation Cycles of Sorbent-Catalyst HART-49